## **IN THE SPECIFICATION:**

At page 54, lines 13-15, please insert a trademark symbol, TM, wherein indicated:

--The amplified cDNA was inserted into pFastBac<sup>TM</sup> donor plasmids (Gibco BRL) at the 3'-end of a honeybee melityin signal peptide and at the 5' end of the Fc sequence of either human IgGl or mouse IgG2b.--

At page 59, lines 19-26, please insert trademark symbols, TM, wherein indicated:

-- In brief, total RNA was purified from each of hybridomas F2-103, F5-77 and F5-157 using Tri-Reagent<sup>TM</sup> according to the manufacturer's instructions (Molecular Research Center, Inc., Cincinnati, Ohio). Full length cDNA was synthesized from total RNA using the SMART RACE<sup>TM</sup> cDNA Amplification Kit (Clontech Laboratories, Inc., Palo Alto, CA) and Superscript II RT<sup>TM</sup> (GibcoBRL). The 5' variable regions of the human heavy and human light chains were isolated by PCR using 5'-RACE<sup>TM</sup> PCR as described by the manufacturer (Clontech Laboratories, Inc.).--

At page 60, lines 5-9, please insert a trademark symbol, TM, wherein indicated:

--Full length PCR products were gel purified and blunt end ligated into SrfI cut PCR-Script<sup>TM</sup> (Stratagene, La Jolla, CA) or PCR-Blunt (Invitrogen, Carlsbad, CA) and sequenced by CFAR, Molecular Biology Core Facility (University of California, San Diego).--

At page 62, lines 37-39, please insert a trademark symbol, TM, wherein indicated:

--Cells were electroporated in a Gene Pulser II (BioRad) set at 240 volts, capacitance=0.950 with a constant time of 15-25 msec.--